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First evidence of a bipartite medial cuneiform in the hominin fossil record: a case report from the Early Pleistocene site of Dmanisi

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Abstract

A medial cuneiform exhibiting complete bipartition was discovered at the Early Pleistocene site of Dmanisi, Georgia. The specimen is the oldest known instance of this anatomical variant in the hominin fossil record. Here we compare developmental variation of the medial cuneiform in fossil hominins, extant humans and great apes, and discuss potential implications of bipartition for hominin foot phylogeny and function. Complete bipartition is rare among modern humans (< 1%); incomplete bipartition was found in 2 of 200 examined great ape specimens and also appears in the form of a divided distal articular surface in the Stw573c Australopithecus africanus specimen. Although various developmental pathways lead to medial cuneiform bipartition, it appears that the bipartite bone does not deviate significantly from normal overall morphology. Together, these data indicate that bipartition represents a phyletically old developmental variant of the medial cuneiform, which does not, however, affect the species-specific morphology and function of this bone.

Key words Dmanisi; morphogenesis; os cuneiforme mediale bipartitum; tarsal bones.

Introduction

The Early Pleistocene site of Dmanisi, Georgia provides a rich assemblage of hominin fossils dated to approximately 1.77 Ma (Vekua et al. 2002; Lordkipanidze et al. 2005, 2006, 2007). During the 2002 excavation season, two nearly complete hominin bones (D4111 and D4112) were discovered in the Block 2 excavation area (square N64 61, layer B1x). Upon subsequent examination, the bones were found to articulate into a single right medial cuneiform. Medial cuneiform bipartition in Dmanisi is similar to what has been described in earlier studies of modern human samples (Barclay, 1932; Barlow, 1942; Kjellström, 2004). The two portions of the Dmanisi bipartite medial cuneiform are referred to subsequently as D4111a/b. D4111a/b is associated anatomically with the first metatarsal D3442 (Lordkipanidze et al. 2007). According to modern human developmental standards, these elements represent an adult individual (the D3442 metatarsal has a fully fused proximal epiphysis; in modern humans, metatarsal I fusion ages are 13–15 years for females and 16–18 years for males, and the medial cuneiform reaches adult morphology by about 6 years of age; Scheuer & Black, 2000).

Bipartition of different human skeletal elements is a relatively frequent subject of comparative anatomical studies. By definition, bipartition means the division of one whole element into two separately-formed parts. Bipartition of bones does not always necessarily equate to two separate, but equal, segments. Bipartition of several carpal and tarsal bones has been described in humans, such as the lunate (Gruber, 1883, 1884; Eggimann, 1951; Schmitt & Schmitt, 1983), scaphoid (Randelli, 1961; Richards et al. 1987) and navicular (Volk, 1937; Zimmer, 1938; Hubner, 1953; Mau, 1960; Wiley & Brown, 1981; Shawdon et al. 1995). The most frequently observed example of a bipartite condition amongst tarsal bones, however, is the medial cuneiform (os cuneiforme mediale bipartitum), first described by Smith (1866) (see Table 1). Bipartition of the medial cuneiform typically results in the bone being divided into upper (dorsal) and lower (plantar) elements. Both of the bipartite elements collectively represent the same structure as appears in non-bipartite medial cuneiforms (Fig. 1).

In the present case study, we provide an anatomical description of a bipartite medial cuneiform from Dmanisi, which is the earliest instance of this anatomical condition that has been reported in the hominin fossil record. We perform morphometric comparisons with five modern human...
populations, including bipartite samples from historical grave burials dated to the 9th–11th centuries in Sigtuna, Sweden (Kjellström, 2004). We also compare the morphology of the Dmanisi medial cuneiform and its associated metatarsal with similar elements in other fossil hominins, focusing on the morphology of the joint between the medial cuneiform and metatarsal I. Using this evidence, we ask whether bipartition of the medial cuneiform may be functionally advantageous during the evolution of hominin locomotion or whether it represents anatomical variation arising from variation in developmental pathways.

### Materials and methods

Data on the incidence and morphological variants of medial cuneiform bipartition in modern human populations and fossil hominins were compiled from the literature. Comparative data on frequencies amongst hominoids were collected from a sample (n = 200) comprised of Pan troglodytes, P. paniscus, Gorilla gorilla and Pongo pygmaeus specimens from the Royal Museum of Central Africa (RMCA), Tervuren and the Anthropological Institute and Museum (AIM), University of Zurich. Data for non-hominoid primate species were compiled from the literature.
The modern human sample of medial cuneiforms used for morphometric comparisons includes data from five populations: recent African-Americans \( (n = 20) \); archaeological specimens from St Sisinius cemetery (11th–12th century), Laas, South Tyrol \( (n = 20) \) (A. H. Schultz Collection, Anthropological Institute, University of Zurich); archaeological specimens from Taforalt, Morocco (ca. 12,000 BP) \( (n = 26) \) (Mariotti et al. 2009) and Afalou bou-Rhumel, Algeria (10,500-8,500 BP) \( (n = 12) \) (Mirazón Lahr & Arensburg, 1995) (Institut de Paléontologie Humaine, Paris); and a sample of bipartite medial cuneiforms from a 9th–11th century mass grave in Sigtuna, Sweden \( (n = 6) \) (Kjellström, 2004).

Linear dimensions of normal and bipartite medial cuneiforms were measured to quantify the overall size and articular dimensions of the bone, following the measurement definitions of Martin & Saller (1957) and Trinkaus (1983) (see Fig. 2). Superior length (SL) is the distance between the most projecting point of the superior (dorsal) edge of the proximal joint area and the corresponding point of the distal joint area of the bone. Middle length (ML) is the distance between the most projecting point of the middle edge of the proximal articular area and the corresponding point of the distal articular area of the bone. Inferior length (IL) is the distance between the most projecting points of the lower (plantar) edge of the proximal articular area and the corresponding point of the distal articular area of the bone. Proximal height (PH) is the distance between the highest (dorsal) point of the superior edge and the deepest (plantar) point of the base of the bone at the proximal end (i.e. not at the edges of the articular area). Proximal articular height (PAH) is the distance between the highest (dorsal) and the deepest (plantar) points of the proximal joint area. Proximal articular breadth (PAB) (Trinkaus, 1983) is the distance between the medial and lateral points at the middle of the proximal joint area, perpendicular to PAH. Distal height is the distance between the highest (dorsal) point of the superior edge and the deepest (plantar) point of the base of the bone at the distal end. Distal articular height (DAH) is the distance between the highest (dorsal) and the deepest (plantar) points of the distal joint area. Distal articular breadth (DAB) (Trinkaus, 1983) is the distance between the medial and the lateral point at the middle of the distal joint area, perpendicular to DAH. All measurements were taken using sliding calipers with a precision of 0.1 mm. Means, SDs and ranges are presented in Table 2. Using these measurements, the following additional variables were calculated in order to estimate articular surface areas: proximal articular area \((PAH \times PAB)\) and distal articular area \((DAH \times DAB)\).

**Results**

**Review of reported cases of bipartite medial cuneiforms**

All reported cases of a bipartite medial cuneiform in modern humans can be assigned to one of three morphological categories (Fig. 1).

1. **Complete bipartition**, in which the medial cuneiform is divided into separate upper (dorsal) and lower (plantar) elements (Fig. 1A, B).
2. **Incomplete bipartition**, in which the two segments are partially fused, with a well-marked proximo-distal cleft on the medial and lateral surfaces demarcating dorsal and plantar segments (Fig. 1C).
Table 2: Linear dimensions of medial cuneiforms (mm).

<table>
<thead>
<tr>
<th>Variables</th>
<th>Afro-American (N = 19)</th>
<th>Laas (N = 20)</th>
<th>Taforalt (N = 26)</th>
<th>Afalou (N = 12)</th>
<th>Sigtuna (N = 6)</th>
<th>Dmanisi (N = 20)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SL</td>
<td>24.87 ± 1.45</td>
<td>23.71 ± 2.26</td>
<td>24.94 ± 1.53</td>
<td>24.87 ± 1.70</td>
<td>24.62 ± 0.82</td>
<td>16.40</td>
</tr>
<tr>
<td>IL</td>
<td>23.80 ± 1.39</td>
<td>21.91 ± 1.58</td>
<td>24.45 ± 1.47</td>
<td>24.17 ± 1.42</td>
<td>22.50 ± 1.36</td>
<td>16.60</td>
</tr>
<tr>
<td>PH</td>
<td>27.51 ± 1.67</td>
<td>25.39 ± 2.18</td>
<td>26.57 ± 1.46</td>
<td>26.57 ± 2.69</td>
<td>27.38 ± 1.10</td>
<td>20.10</td>
</tr>
<tr>
<td>PAB</td>
<td>23.11 ± 1.56</td>
<td>21.32 ± 1.95</td>
<td>20.29 ± 2.10</td>
<td>20.80 ± 1.91</td>
<td>24.57 ± 3.47</td>
<td>13.10</td>
</tr>
<tr>
<td>PAB</td>
<td>15.99 ± 1.36</td>
<td>15.86 ± 1.65</td>
<td>15.81 ± 1.36</td>
<td>15.93 ± 1.39</td>
<td>16.58 ± 0.75</td>
<td>13.50</td>
</tr>
<tr>
<td>DH</td>
<td>33.84 ± 2.51</td>
<td>30.96 ± 2.29</td>
<td>32.80 ± 1.94</td>
<td>33.18 ± 2.40</td>
<td>34.08 ± 1.51</td>
<td>28.80</td>
</tr>
<tr>
<td>DH</td>
<td>30.05–38.50</td>
<td>27.70–34.90</td>
<td>29.43–35.90</td>
<td>27.30–35.86</td>
<td>32.10–35.90</td>
<td></td>
</tr>
<tr>
<td>DAH</td>
<td>30.88 ± 2.07</td>
<td>28.17 ± 1.85</td>
<td>29.38 ± 1.74</td>
<td>29.50 ± 2.30</td>
<td>30.45 ± 1.20</td>
<td>26.40</td>
</tr>
<tr>
<td>DAB</td>
<td>13.14 ± 1.20</td>
<td>13.03 ± 2.15</td>
<td>12.63 ± 1.51</td>
<td>13.64 ± 1.18</td>
<td>13.80 ± 0.88</td>
<td>12.50</td>
</tr>
</tbody>
</table>

SL, superior length; ML, middle length; IL, inferior length; PH, proximal height; PAH, proximal articular height; PAB, proximal articular breadth; DH, distal height; DAH, distal articular height; DAB, distal articular breadth.

Data represent mean ± SD (first row), and range (second row).

3 Division of the distal articular surface (Fig. 1D). The latter variant seems to be the most commonly observed (Barlow, 1942).

Bipartition can affect one or both feet of an individual, which is referred to here as unilateral and bilateral cases, respectively. Table 1 provides a list of reported cases of each category, demonstrating that bipartition is relatively rare. In the largest sample examined so far, an incidence of 0.27 and 0.39% was reported for complete and partial bipartitions, respectively (Gruber, 1877). A high frequency of bipartition has been encountered in populations of low genetic diversity, such as bilateral bipartition to various degrees in three individuals from a burial site in Sigtuna, Sweden (Kjellström, 2004). This indicates a heritable component of the trait (Gruber, 1884; Pfitzner, 1896; Barlow, 1942; Azurza & Sakellariou, 2001; Kjellström, 2004). Although a majority of reports of bipartite medial cuneiforms come from archaeological, osteological or dissection material, the wide use of magnetic resonance imaging and computed tomography in clinical diagnostics of midfoot trauma and pathology has yielded a growing number of cases as incidental clinical findings.¹

In the bipartite medial cuneiform, an extra-articular joint divides the bone into plantar and dorsal segments. The size relationship between these elements is variable and appears to be case-specific. Most commonly, there are relatively large plantar and relatively small dorsal portions. Some authors describe the synovial joint as plane-like, similar to joints between other tarsal bones that permit gliding movements (Barlow, 1942). Others propose a cartilaginous synchondrosis (Azurza & Sakellariou, 2001), where the hyaline cartilage (a relic of the embryonic cartilage matrix of the skeleton) creates an immobile synarthrosis joint. In either case, the volume of a bipartite medial cuneiform is slightly larger than that of a typical non-bipartite medial cuneiform. Muscles attaching to a bipartite medial cuneiform are only received by the plantar segment. A slip of the tendons of tibialis posterior and peroneus longus typically attaches to the plantar surface, whereas the tibialis anterior tendon typically attaches to the medial surface of the plantar segment. No muscles are attached to the lateral area of the dorsal segment but instead there is an attachment of an interosseous ligament (Barlow, 1942). Descriptions of first metatarsals associated with bipartite medial cuneiforms (all categories) report expansion of the

¹In all of these cases, medial cuneiform bipartition is asymptomatic and not related to the diagnosed disorder (Dellacorte et al. 1992; Azurza and Sakellariou, 2001; Bismil et al. 2005; Elias et al. 2008).
dorso-plantar diameter and division of the proximal articular surface into two facets separated by a ridge. The upper facet usually is concave, articulating with the anterior facet of the dorsal segment of the bipartite medial cuneiform, whereas the plantar facet usually is slightly convex, articulating with the plantar cuneiform segment (Table 1).

Review of reported cases of bipartite medial cuneiforms in non-human primates

Bipartition was observed in two of 200 hominoid specimens. One instance of bipartition was observed in a 6-year-old *Pongo pygmaeus abelii* (Zurich collection id: AIM10141; Fig. 3A–C), whereas the other instance was observed in an adult *P. paniscus* (Tevuren collection id: RMCA15296; Fig. 3D–F). Both of these individuals were classified as incomplete (category 2) bipartition, where a well-marked proximo-distal cleft on the medial and lateral surfaces divides the bone into two regions, dorsal and plantar. These are the first cases of bipartition reported in non-human primates.

Review of medial cuneiforms in Plio-Pleistocene hominin fossils

Several medial cuneiforms have been described in the hominin Plio-Pleistocene fossil record: a left, complete one, possibly the earliest available *Australopithecus africanus* specimen (Stw573c) (Clarke & Tobias, 1995); a right, adult *Australopithecus afarensis* specimen (AL333-28) conserving the plantar two-thirds of the element (Latimer et al. 1982); a left, fragmentary specimen attributed to either *Paranthropus robustus* or *Homo cf. erectus* (SKX31117) (Susman, 1989)\(^2\); and a left, complete *H. habilis* specimen (OH8e) (Day & Napier, 1964). Medial cuneiforms have also been reported for *H. floresiensis* (Jungers et al. 2009) and *Ardipithecus ramidus* (Lovejoy et al. 2009) but detailed descriptions of their morphology are not yet available.

In initial studies of the Stw573c medial cuneiform (Clarke & Tobias, 1995; Deloison, 2004), the distal articular facet was described as extending onto the medial surface and having a ‘less square-shaped’ configuration. Such morphology resembles *Pan* (Deloison, 2004). The distal facet of Stw573c is markedly convex. The L-shaped facet for the intermediate cuneiform resembles the condition in modern humans. A recent quantitative analysis (McHenry & Jones, 2006), however, demonstrated that the distal articular facet does not extend onto the medial surface and that the medial cuneiform-metatarsal facet is human-like. A photograph of the distal aspect of this specimen (Deloison, 2004) suggests the presence of two separate distal facets instead of one. Therefore, it is likely that the Stw573c medial cuneiform has a divided distal articular surface (category 3), whereas the proximal surface has a single, undivided facet. In AL333-28, the preserved two-thirds of the distal articulation show a convex facet with a distinct longitudinal-dorso-plantar ridge (Latimer et al. 1982), similar to what has been described in Stw573c. Re-examination of this surface shows no signs of bipartition. Rather, the apparent ridges are due to partial erosion of the articular surface (Tim White, Department of Integrative Biology, University of California, Berkeley, and Berhane Asfaw, Rift Valley Research Service, Addis Ababa, pers. comm.)

The OH8e and SKX31117 medial cuneiforms are different from those of the two australopithecine specimens, primarily in that the former two have a less convex distal articular surface (Day & Napier, 1964; Susman, 1989).

The first metatarsal (Stw573d), which is associated with the Stw573c medial cuneiform, has a slightly concave proximal articular surface divided into upper and lower facets by a transverse ridge (Latimer et al. 1982; Clarke & Tobias, 1995). A first left metatarsal AL333-54, which was found in the same archaeological layer as the AL333-28 medial cuneiform, was described as having a similar morphology. Contrastingly, in the OH8 foot, the first metatarsal OH8h, which articulates with the medial cuneiform OH8e, has a flat proximal articular surface and a similar morphology has been described for the first metatarsal SKX5017, which corresponds in size to the SKX31117 medial cuneiform (Susman, 1989).

\(^2\)Re-examination of the SKX31117 medial cuneiform by one of the authors (TJ) indicates that this specimen does not represent a medial cuneiform and is probably non-hominin. Future investigation is needed to clarify this issue. The description used here follows Susman (1989).
The D4111a/b bipartite medial cuneiform and associated metatarsal D3442

D4111a/b is represented by a complete, undistorted and well-preserved pair of segments (Fig. 4). Post-mortem abrasions are present on surfaces of both segments. The plantar segment has only minor areas of abrasion on the lateral side (Fig. 4B′), which continue onto the articular surface of the dorsal part. The dorsal segment also has small abrasions on the lateral side, as well as on the articular surface for the base of the second metatarsal (Fig. 4B,F). The dorsal segment also exhibits a small amount of damage on the medial edge of its articular surface with the plantar segment (Fig. 4C).

The dorsal segment is relatively small and ovoid in shape (Fig. 4A–F), whereas the plantar segment is larger anteroposteriorly and cylindrical in shape (Fig. 4A′–F′). Articulation of the two segments creates a well-defined joint surface; the plantar surface of the dorsal segment is slightly convex (Fig. 4F), whereas the dorsal surface of the plantar segment is slightly concave (Fig. 4E′). The articulation between the two segments has a similarly smooth structure as the other articular surfaces on the bone, probably indicating the presence of a plane synovial joint covered by cartilage and synovial membrane.

The dorsal segment has a flattened articular surface for the base of the first metatarsal on its dorsal aspect, similar to what is reported in modern humans and the OH8e specimen (Fig. 4D). On the proximal side of the dorsal segment, there is no articular surface for the navicular, as has been observed in cases of bipartite medial cuneiforms in modern humans (Barlow, 1942; Kjellström, 2004). Medial and lateral surfaces join each other at a ridge that demarcates the articular surface for the intermediate cuneiform and a roughened surface with well-defined tubercles for insertion of dorsal cuneonavicular ligaments proximally and the dorsal ligament of the first tarsometatarsal joint distally. The articular surface for the intermediate cuneiform is large and ovoid in form, and flattened with a dorso-plantar inclination on its lateral aspect (Fig. 4C). This form of articulation differs from the more narrow and elongated form that has been described in modern human samples, Stw573c and OH8e. The lateral side of the dorsal segment continues distally towards the articulation for the base of the second metatarsal (Fig. 4B). This surface, along with the articular surface for the intermediate cuneiform, forms a longitudinal ridge that is more closed-angled compared with modern humans and Stw573c/OH8e.

The plantar segment is divided distally by a longitudinal cleft forming a convexity on the articular surface for the base of the first metatarsal. Similar morphology is reported for the australopithecine specimens Stw573c and AL333-28 (Latimer et al. 1982; Clarke & Tobias, 1995); this morphology differs from the flat articular facet in modern humans and OH8e (Fig. 4D′). Although proximally the articular surface for the navicular is concave like in modern humans and Plio-Pleistocene hominins, it has an ovoid form that is elon-
gated dorso-medially to planto-laterally and facing laterally unlike in modern humans and Plio-Pleistocene hominins (Fig. 4C). The medial surface of the plantar segment has a prominent tubercle for insertion of the tendon of \textit{tibialis anterior} on its most anteroinferior angle (Fig. 4A). The plantar surface of the plantar segment has a well-defined tubercle for the insertion of a slip of the tendon of \textit{tibialis posterior} on its proximal side and, on its distal side, a tubercle for the insertion of the \textit{peroneus longus} tendon (Fig. 4F).

The D3442 right first metatarsal (Fig. 4G–I) is large in dorso-plantar diameter compared with a typical modern human first metatarsal. It has a proximal articular surface divided into two facets by a horizontal ridge. The dorsal facet is mediolaterally concave, articulating with the anterior facet of the dorsal part of the medial cuneiform. The plantar facet is concave, articulating with the anterior convex facet of the plantar part of the medial cuneiform. Together, the two facets form an ‘8-shaped’ articular contour, similar to the morphology reported for Stw753d and AL333-54. An ‘8-shaped’ articular contour of the proximal first metatarsal has a low frequency in modern humans (2%) (Hartman & Mordret, 1889) and this condition is first metatarsal has a low frequency in modern humans; (ii) size-related (allometric) differences; and (iii) differences between normal and bipartite medial cuneiforms.

Figure 5 shows that the bipartite cuneiform sample from Sigtuna is well within the mode of shape variation of human non-bipartite medial cuneiforms. Dmanisi is clearly separated from modern humans and differences in shape cannot be explained by differences in size alone. It is thus sensible to assume that differences between Dmanisi and the modern human sample do not reflect differences in the bipartite condition or allometric scaling but taxon-specific differences. The Dmanisi medial cuneiform is characterized by a relatively large inferior length and a relatively large distal articular surface, in congruence with the large base of metatarsal I.

### Discussion

**Evolutionary developmental origins of the hominin foot and of medial cuneiform bipartition**

Elements of the autopodium first appear in Devonian polydactylous tetrapods with six to eight digits (Coates & Clack, 1990). The pattern of basic tarsal units has undergone various evolutionary variations, including fusion or loss of elements and evolution of entirely new elements (Schaeffer, 1941). Because of the lack of evidence for successive fossil anatomy, it is difficult to infer homology within structures (Lewis, 1989). Differential fusion of the tarsal elements produces a wide spectrum of evolutionary developmental variation. In some species, separate cartilaginous precursors, which join during development to form one bone, bear evidence of phylogenetic history, whereas in other species, tarsal elements are highly derived and do not permit phylogenetic inferences (Lewis, 1989). Figure 6 shows the hypothetical basic tarsal units represented in human foot primordia; these include the talus (homologous to \textit{os tibiale intermedium} and \textit{os tibiale centrale proximale}), calcaneus (\textit{os fibulare} and \textit{os pisiforme}), navicular (\textit{tibiale centrale distale} and \textit{fibulare centrale distale}), tuberosity of the navicular (\textit{os tibiale externum tarsi}), medial cuneiform (\textit{os tarsale distale I} and distal end of the prehallux primordium),

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**Fig. 5** Medial cuneiform shape variation. Plot of ratio between distal and proximal articular surface areas (DAA : proximal articular area (PAA)) vs. ratio between inferior and superior lengths (IL : SL). Black/white circles, Taforalt/Af Aliou; black/white squares, Laas/Afro-American; asterisks, Sigtuna bipartite medial cuneiforms; X, Dmanisi D4111.
intermediate cuneiform (os tarsale distale II), lateral cuneiform (os tarsale distale III) and cuboid (os tarsale distale IV) (Čihák, 1972; Berman & Henrici, 2003).

Figures 7 and 8 give an overview of documented developmental pathways of the human medial cuneiform. Most frequently, it originates from a single mesenchymal primordium that ultimately develops into a single complete bone (Figs 7B and 8b1) (Scheuer & Black, 2000). In rare cases, a single complete medial cuneiform can arise from divided plantar and dorsal primordia that are separated by non-chondrified tissue (Figs 7A and 8b2) (Gardner et al. 1959; Čihák, 1972). The non-chondrified tissue constitutes a closely-packed mesenchymal-looking, cell-forming interzone (Pacifici et al. 2005) that eventually disappears and is replaced by other proliferating cells giving rise to a single complete bone (Fig. 8c1 e1). Fusion of the two transient primordia (Figs 7A and 8b2 e1) is interpreted as fusion of the os tarsale distale I and the prehallucial primordium (Čihák, 1972).

Bipartition can arise along two different developmental pathways (Barlow, 1942). Initial division in the mesenchymal primordium gives rise to a joint between two ossification centers (Figs 7A and 8b2 c2), resulting in a bipartite medial cuneiform (Fig. 8c2 e2) (Čihák, 1972). Alternatively, a single cartilaginous anlage gives rise to two ossification centers with a small fibrous cleft representing a hyaline spondylodesis (growth plate) between them (Figs 7B and 8c1 d2), eventually resulting in two articulating bones (Fig. 8d2 e2) (Pfitzner, 1896; O’Neal et al. 1995).

Genetic factors underlying variation in medial cuneiform developmental pathways might be identified at two different levels, namely Hox and Sox transcription factors, which specify the patterning and shape of the embryonic skeleton (Hall & Miyake, 2002; Cohen, 2006; Montero & Hurlé, 2007), or local regulators of mesenchymal condensation, which control the shape of the skeletal elements via autocrine/paracrine factors (Garcidiego-Cazares et al. 2004; Hentschel et al. 2004; Pacifici et al. 2005; Cohen, 2006; Newman & Bhat, 2007). In the early stages of development, formation of a divided mesenchymal primordium (Fig. 8b2) could result from altered cell-intrinsic patterning at the gene level and/or variation in activator/inhibitor positional signaling mediators responsible for local control of the
mesenchymal primordium. In later stages of development, formation of two centers of ossification inside one single cartilaginous anlage (Fig. 8d2) could result from variation in activator/inhibitor positional signaling mediators responsible for local control of the cartilaginous anlage (Newman & Bhat, 2007). By the end of the adult stage of development, the outcome of both developmental scenarios would be a bipartite bone (Fig. 8e2), unless the separate bone precursors fuse producing a complete bone (Fig. 8e1).

Modern humans exhibit several variations in tarsal bone expression at low frequencies (Marti, 1947). These reflect homologous variants to the examples mentioned earlier: os trigonum (Zeichen et al. 1999; Chao, 2004; Mouhsine et al. 2004), os perineum (cuboideum secundarium, centrale 4) (Bloom, 1991), os sustentaculi (Bloom et al. 1986), os naviculare bipartitum (Cotta, 1961) and os vesalianum (Virchow, 1922; Inoue et al. 1999; Boya et al. 2005). A supernumerary ossicle (‘prehallux’) situated at the disto-medial border of the medial cuneiform, or proximo-medial border of the first metatarsal, is common in Ceboidea and Hylobatidae. Such an ossicle also has been observed in pongids and Homo (Lewis, 1972; Wikander et al. 1986). The ‘prehallux’ may be a relict skeletal element of a pre-axial fin ray from the earliest phylogenetic stages of vertebrates (Lewis, 1972).

These non-random developmental variants provide the raw material on which selection has the potential to act during evolution (Alberch, 1983; Erlebacher et al. 1995). Morphological variants of the medial cuneiform also seem to result from a non-random pattern of developmental variation, i.e. different degrees of bipartition. Regarding the evolutionary development origins of bipartition, it may be speculated that the medial cuneiform is homologous to os tarsale distale I and the distal part of the prehallux primordium (Čihák, 1972; Berman & Henrič, 2003).

Structure and function
In all clinical reports of a bipartite medial cuneiform, the condition was discovered incidentally during evaluation of subjects for a foot injury or a degenerative process in articular surfaces rather than an association with functional limitations of the foot (Dellacorte et al. 1992; O’Neal et al. 1995; Sener, 1999; Azurza & Sakellariou, 2001; Chiodo et al. 2002; Bismil et al. 2005; Fulwadhva & Parker, 2007). Presumably, the relatively flattened planar joint between the halves of the bipartite medial cuneiform does not restrict or alter motion of the medial ray of the foot.

In our modern human comparative sample, there was no indication that the bipartite medial cuneiforms of the Sigtuna sample significantly exceeded normal patterns of cuneiform shape variation. Morphological differences between the Dmanisi bipartite medial cuneiform and modern human cuneiforms thus most likely reflect species-specific differences, irrespective of whether these bones display bipartition. The articulation between the two segments of the Dmanisi medial cuneiform has a smooth structure, similar to the structure of other articular surfaces on these bones. This pattern is also apparent in articular surfaces of modern human bipartite medial cuneiforms. It is thus probable that the intracuneiform joint of Dmanisi, as in modern humans, was covered by cartilage and a synovial membrane.

Overall, it appears that the bipartite condition in the medial cuneiform represents developmental variation that does not cause significant overall morphological differences. Presumably the lack of morphological differences also implies a lack of functional differences. This provides an interesting perspective on the relationship of developmental and adaptive/functional constraints. The fact that the taxon-specific (most likely functionally relevant) morphology of the medial cuneiform can be reached by
different developmental pathways, some of which imply bipartition to various degrees, points toward higher-order, epigenetic constraints that canalize the development of midfoot morphology as a whole. This indicates morphogenetic homeostasis in the sense that foot ontogeny could be buffered against environmental noise as well as against developmental noise. Accordingly, it appears that the network of developmental pathways graphed in Fig. 8 not only gives rise to patterns of medial cuneiform variation but also provides the required developmental homeostasis, as developmental disturbance at any node or link in the network can be compensated by alternative pathways.

Fossil hominin medial cuneiforms and first metatarsals

In hominoids, the distal articular surface of the medial cuneiform is convex, wide and medially-oriented, such that the articulating hallux is medially divergent. In humans this surface is flat, narrow and anteriorly-facing, such that the hallux does not diverge medially (Lewis, 1980, 1989). The latter condition exists in all described fossil hominin medial cuneiforms, which indicates that none of them are likely to have had a divergent hallux (McHenry & Jones, 2006). The Dmanisi bipartite medial cuneiform parallels this pattern.

In modern humans, morphology of the proximal joint surface of the first metatarsal (i.e. the surface articulating with the cuneiform) typically reflects the distal joint surface of the medial cuneiform. It is flat in the case of a single, non-bipartite medial cuneiform and it displays two facets separated by a transverse ridge (‘B-shaped’ circumference) in the case of a bipartite medial cuneiform. The latter association is observed in the relatively convex/concave first tarsometatarsal joint of the Dmanisi D3442/D4111a/b bones.

The situation is more complex in other Plio-Pleistocene hominins (Harcourt-Smith & Aiello, 2004; DeSilva, 2008; Proctor et al. 2008). The first tarsometatarsal joint surface is flat and undivided in OH8 and in SKX5017/SKX31117 (both attributed to Homo). In the Stw573 foot (attributed to Australopithecus), the proximal metatarsal surface has two facets. There is evidence that the same is true for the distal surface of the medial cuneiform. In the A. afarensis first metatarsal (AL333-54), the proximal articular surface exhibits two facets, whereas the only partially preserved medial cuneiform from the same stratigraphic location (AL333-28) bears no evidence of bipartition. However, not all first metatarsals attributed to Australopithecus reportedly exhibit a double facet (Day & Napier, 1964; Susman, 1989).

If we assume that a double-faceted proximal metatarsal joint surface is indicative of bipartition of the medial cuneiform in any of the three categories, it appears that this condition was relatively frequent in Plio-Pleistocene hominins compared with modern human populations. The significance of a potentially higher incidence remains to be clarified. It could reflect increased developmental variation during evolutionary diversification of the hominin foot but it could also represent a sampling artifact.

Conclusions

The Dmanisi D4111a/b bipartite medial cuneiform is the oldest known instance of this condition in the hominin fossil record. Incomplete bipartition in the form of a divided distal articular surface also appears in the medial cuneiform of the A. africanaus foot Stw573, although this is not definite. In modern humans, bipartition has been reported to be rare and we find similarly low incidences (incomplete bipartition) in large samples of Pan and Pongo. Bipartition of the medial cuneiform is associated with a divided proximal articular surface of the first metatarsal in modern humans, in D3442/D4111a/b, and possibly in the Stw573 foot. Accordingly, the isolated A. afarensis metatarsal AL333-54, which exhibits a divided proximal surface, might also have been associated with a bipartite medial cuneiform. This raises the question whether this condition was more frequent in Plio-Pleistocene hominins than in extant humans and great apes.

In all linear dimensions, the Dmanisi bipartite medial cuneiform falls below the corresponding mean values for medial cuneiforms of modern human populations. In shape, the Dmanisi specimen differs significantly from modern human medial cuneiforms, indicating species-specific morphologies irrespective of the presence or absence of bipartition.

Various developmental pathways lead to bipartition or the normal condition of the medial cuneiform. Although these pathways generate developmental variation, the overall shape of the bone remains conserved. This indicates that the morphology of the medial cuneiform is constrained by higher-order processes of developmental integration. It is thus likely that bipartition is of no functional relevance.

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Bipartite medial cuneiform from Dmanisi, T. Jashashvili et al.


